



(19)

Europäisches Patentamt

European Patent Office

Office européen des brevets



(11)

EP 0 794 007 A1

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication:  
10.09.1997 Bulletin 1997/37

(51) Int. Cl.<sup>6</sup>: B01L 3/14, B32B 27/00,  
C23C 14/00

(21) Application number: 96103454.3

(22) Date of filing: 06.03.1996

(84) Designated Contracting States:  
DE ES FR GB IT

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(54) Blood collection tube assembly

(57) The present invention is a plastic container coated with a multi-layer barrier coating. The multi-layer barrier coating is useful for providing an effective barrier against gas permeability in containers and for extending shelf-life of containers, especially plastic evacuated blood collection devices.

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**Description****BACKGROUND OF THE INVENTION****5 1. Field of the Invention**

This invention relates to a multi-layer barrier coating for providing an effective barrier against gas and water permeability for containers, especially plastic blood collection tubes.

**10 2. Description of the Related Art**

With the increased emphasis on the use of plastic medical products, a special need exists for improving the barrier properties of articles made of polymers.

Such medical products that would derive a considerable benefit from improving their barrier properties include, but 15 are not limited to, collection tubes and particularly those used for blood collection.

Blood collection tubes require certain performance standards to be acceptable for use in medical applications. Such performance standards include the ability to maintain greater than about 90% original draw volume over a one year period, to be radiation sterilizable and to be non-interfering in tests and analysis.

Therefore, a need exists to improve the barrier properties of articles made of polymers and in particular plastic 20 evacuated blood collection tubes wherein certain performance standards would be met and the article would be effective and usable in medical applications.

**SUMMARY OF THE INVENTION**

25 The present invention is a plastic composite container with a multi-layer barrier coating comprising at least two barrier materials disposed over the outer surface of the previously formed composite container. Desirably, the barrier materials comprise a first layer of a polymeric material and a second layer of an inorganic material applied over the first layer.

The first layer, a primer coating, is preferably a highly cross linked acrylate polymer. The coating may be formed either on an interior surface portion, on an exterior surface portion, or both of the container.

30 The second layer of the barrier coating may preferably be a silicon oxide based composition, such as  $\text{SiO}_x$  wherein  $x$  is from 1.0 to about 2.0; or an aluminium oxide based composition. Most preferably, the second layer is a silicon oxide based composition applied over the first layer.

Preferably, the primer coating is a blend of monoacrylate (i.e., isobornyl acrylate) and diacrylate monomers (i.e., an 35 epoxy diacrylate or a urethane diacrylate) as described in U.S. Patent Nos. 4,490,774, 4,696,719, 4,647,818, 4,842,893, 4,954,371 and 5,032,461, the disclosures of which are herein incorporated by reference. The primer coating is cured by an electron beam or by a source of ultraviolet radiation.

Desirably, the first layer is formed of a substantially cross linked component selected from the group consisting of 40 polyacrylates and mixtures of polyacrylates and monacrylates having an average molecular weight of between 150 and 1,000 and a vapor pressure in the range of  $1 \times 10^{-6}$  to  $1 \times 10^{-1}$  Torr at standard temperature and pressure. Most preferably, the material is a diacrylate.

The process for applying the primer coating to a container is preferably carried out in a vacuum chamber wherein 45 a curable monomer component is metered to a heated vaporizer system where the material is atomized, vaporized and condensed on the surface of the container. Following deposit of the monomer onto the surface of the container, it is cured by suitable means such as electron beam curing. The deposition and curing steps may be repeated until the desired number of layers has been achieved.

Preferably, the thickness of the acrylate primer coating is about .1 to about 10 microns and most preferably from about .5 to about 5 microns.

A desirable second layer which is disposed over the first layer preferably comprises a silicon oxide based composition, such as  $\text{SiO}_x$ . Such a film desirably is derived from volatile organosilicon compounds.

50 A method for depositing a silicon oxide based film is as follows: (a) pretreating the first layer on the container with a first plasma coating of oxygen; (b) controllably flowing a gas stream including an organosilicon compound into a plasma; and (c) depositing a silicon oxide onto the first layer while maintaining a pressure of less than about 500 microns  $\text{H}_2$  during the depositing.

It is believed that the pretreatment step provides for improved adherence qualities between the second layer and 55 the primer coating.

The organosilicon compound is preferably combined with oxygen and optionally helium or another inert gas such as argon or nitrogen and at least a portion of the plasma is preferably magnetically confined adjacent to the surface of the first layer during the depositing, most preferably by an unbalanced magnetron.

The silicon oxide based composition provides a dense, vapor-impervious coating over the primer coating. Prefera-

bly, the thickness of the silicon oxide based layer is about 100 to about 10,000 Angstroms ( $\text{\AA}$ ) and most preferably from about 1,000 to about 3,000  $\text{\AA}$ . A coating above 10,000  $\text{\AA}$  may crack and therefore be ineffective as a barrier.

Plastic tubes coated with the multi-layer barrier coating, comprising the primer coating and an oxide layer are able to maintain substantially far better vacuum retention and draw volume retention than previous tubes comprised of polymer compositions and blends thereof without a coating of barrier materials or of tubes comprising only an oxide coating. In addition, the tube's resistance to impact is much better than that of glass. Most notably is the clarity of the multi-layer coating and its durability to substantially withstand resistance to impact.

5 Most preferably, the container of the present invention is a blood collection device. The blood collection device can be either an evacuated blood collection tube or a non-evacuated blood collection tube. The blood collection tube is  
10 desirably made of polyethylene terephthalate, polypropylene, polyethylene naphthalate or copolymers thereof.

15 Printing may be placed on the multi-layer barrier coating applied to the container of interest. For example, a product identification, bar code, brand name, company logo, lot number, expiration date and other data and information may all be included on the barrier coating. Moreover, a matte finish or a corona discharged surface may be developed on the barrier coating so as to make the surface appropriate for writing additional information on the label. Furthermore, a pressure sensitive adhesive label may be placed over the barrier coating so as to accommodate various hospital over-labels, for example.

Preferably, the multi-layer barrier coating of the present invention provides a transparent or colorless appearance and may have printed matter applied thereon.

20 A further advantage is that the method of the present invention provides a reduction in the gas permeability of three-dimensional objects that has not been achieved with conventional deposition method typically used with thin films.

Surprisingly, it has been found that a primer coating of acrylate on a plastic surface followed by a coating of  $\text{SiO}_x$  will result in an oxide coating containing reduced amounts of elemental carbon, thereby resulting in a better barrier coating as compared to  $\text{SiO}_x$  coated directly over the plastic. It is believed that it is difficult to prepare a uniform continuous gas-impervious coating of  $\text{SiO}_x$  with good adhesion properties and minimal defects on plastic surfaces. It is believed, 25 that poor adhesion properties are due to the presence of low molecular weight hydrocarbon moieties at the surface of the plastic, such as dust particles and imperfections, formed during injection molding. The amorphous regions at the surface of the polyolefin, having high chain mobility, also cause low adhesion of  $\text{SiO}_x$  to a polymer. Such plastic surfaces include polyolefins such as polypropylene and polyethylene.

It has been found that a highly crosslinked layer of acrylate improves the adhesion between a plastic surface and 30  $\text{SiO}_x$  and overall improves the thermomechanical stability of the coated system. In addition, acrylate primer coating has a role of a planarization (leveling) layer, covering the particles and imperfections on the surface of a polymer and reducing the defect density in the deposited inorganic coatings. The good bonding properties of the acrylate are also due to the fact that acrylate is polar and the polar property provides means for good bond formation between the  $\text{SiO}_x$  and the acrylate. In addition, it has been found that a good bond formation is made between plastic tubes made of polypropylene and acrylate. Thus, the present invention provides the means of substantially improving the barrier properties of 35 polypropylene tubes. The adhesion properties of both the acrylate coating and the oxide coating can be further substantially improved by surface pretreatment methods such as flame or oxygen plasma. Therefore, a significant reduction in permeability of the article is due to the substantially improved  $\text{SiO}_x$  surface coverage that is obtained by the use of a primer coating of acrylate on the plastic article surface.

40 A plastic blood collection tube coated with the multi-layer barrier coating of the present invention will not interfere with testing and analysis that is typically performed on blood in a tube. Such tests include but are not limited to, routine chemical analysis, biological inertness, hematology, blood chemistry, blood typing, toxicology analysis or therapeutic drug monitoring and other clinical tests involving body fluids. Furthermore, a plastic blood collection tube coated with the barrier coating is capable of being subjected to automated machinery such as centrifuges and may be exposed to 45 certain levels of radiation in the sterilization process with substantially no change in optical or mechanical and functional properties.

#### DESCRIPTION OF THE DRAWINGS

50 FIG. 1 is a perspective view of a typical blood collection tube with a stopper.  
FIG. 2 is a longitudinal sectional view of the tube of FIG. 1 taken along line 2-2.  
FIG. 3 is a longitudinal sectional view of a tube-shaped container similar to the tube of FIG. 1 without a stopper, comprising a multi-layer barrier coating.

55 FIG. 4 is a longitudinal sectional view of a tube-shaped container, similar to the tube of FIG. 1 with a stopper, comprising a multi-layer barrier coating.

FIG. 5 is a longitudinal sectional view of a further embodiment of the invention illustrating the tube with a stopper similar to FIG. 1 and with the multi-layer barrier coating encompassing both the tube and stopper thereof.

FIG. 6 illustrates an enlarged partially sectioned, diagram of a flash evaporator apparatus.

FIG. 7 illustrates a plasma deposition system.

FIG. 8 illustrates atom percent versus etch time of polypropylene plaque, acrylate and SiO<sub>x</sub>.  
 FIG. 9 illustrates atom percent versus etch time of polypropylene plaque and SiO<sub>x</sub>.

#### DETAILED DESCRIPTION

5 The present invention may be embodied in other specific forms and is not limited to any specific embodiment described in detail which is merely exemplary. Various other modifications will be apparent to and readily made by those skilled in the art without departing from the scope and spirit of the invention. The scope of the invention will be measured by the appended claims and their equivalents.

10 Referring to the drawings in which like reference characters refer to like parts throughout the several views thereof, FIGS. 1 and 2 show a typical blood collection tube 10, having a sidewall 11 extending from an open end 16 to a closed end 18 and a stopper 14 which includes a lower annular portion or skirt 15 which extends into and presses against the inner surface 12 of the sidewall for maintaining stopper 14 in place.

15 FIG. 2 schematically illustrates that there are three mechanisms for a change in vacuum in a blood collection tube: (A) gas permeation through the stopper material; (B) gas permeation through the tube and (C) leak at the stopper tube interface. Therefore, when there is substantially no gas permeation and no leak, there is good vacuum retention and good draw volume retention.

20 FIG. 3 shows the preferred embodiment of the invention, a plastic tube coated with at least two layers of barrier materials. The preferred embodiment includes many components which are substantially identical to the components of FIGS. 1 and 2. Accordingly, similar components performing similar functions will be numbered identically to those components of FIGS. 1 and 2, except that a suffix "a" will be used to identify those components in FIG. 3.

25 Referring now to FIG. 3, the preferred embodiment of the invention, collection tube assembly 20 comprises a plastic tube 10a, having a sidewall 11a extending from an opened end 16a to a closed end 18a. A barrier coating 25 extends over a substantial portion of the outer surface of the tube with the exception of open end 16a. Barrier coating 25 comprises a first layer 26 of an acrylate polymer material and a second layer 27 of a silicon oxide based composition.

30 FIG. 4 illustrates an alternate embodiment of the invention, wherein collection tube assembly 40 comprises stopper 48 in place for closing open end 41 of tube 42. As can be seen, sidewall 43 extends from open end 41 to closed end 44 and stopper 48 includes an annular upper portion 50 which extends over the top edge of tube 42. Stopper 48 includes a lower annular portion or skirt 49 which extends into and presses against the inside inner surface 46 of sidewall 43 for maintaining stopper 48 in place. Also, the stopper has a septum portion 52 for receiving a cannula therethrough.

35 Thus, the user, once receiving a container such as that shown in FIG. 4 with a sample contained therein, may insert a cannula through septum 52 for receiving part or all of the contents in tube 42 to perform various tests on a sample. Covering a substantial portion of the length of the tube is a multi-layer barrier coating 45. Multi-layer barrier coating 45 covers substantially most of the tube with the exception of open end 41 thereof. Multi-layer barrier coating 45 comprises a first layer 54 of a polymer material and a second layer 56 of an inorganic material. FIG. 4 differs from the embodiment in FIG. 3 in that the tube may be evacuated with the simultaneous placement of stopper 48 therein after the application of layers 54 and 56 over the tube. Alternatively, the multi-layer barrier coating may be applied to the tube after it has been evacuated.

40 FIG. 5 shows an additional embodiment of the barrier coating and a tube. The alternate embodiment functions in a similar manner to the embodiment illustrated in FIG. 4. Accordingly, similar components performing similar functions will be numbered identically to those components in the embodiment of FIG. 4, except that a suffix "a" will be used to identify those components in FIG. 5.

45 Referring now to FIG. 5, a further embodiment 60 of the invention wherein multi-layer barrier coating 45a incorporates both upper portion 50a of stopper 48a, as well as the entire outer surface of tube 42a. Multi-layer barrier coating 45a includes serrations 62 at the tube, stopper interface. The serrations are registered so that it can be determined if the sealed container has been tampered with. Such an embodiment may be utilized, for example, for sealing the container with the stopper in place. Once a sample has been placed in the tube, the sample cannot be tampered with by removal of the stopper. Additionally, the serrations may be registered so that it can be determined if the sealed container has been tampered with. Such an arrangement may be appropriate, for example, in drug abuse testing, specimen identification and quality control.

50 In an alternate embodiment of the invention, multi-layer barrier coating 45 is repeatedly applied to the inner and/or outer surface of the tube. Preferably, the coating is applied at least twice.

It will be understood by practitioners-in-the-art, that such tubes may contain reagents in the form of additives or coatings on the inner wall of the tube.

55 The multi-layer barrier coating forms a substantially clear or translucent barrier. Therefore, the contents of a plastic tube with a multi-layer barrier coating comprising at least two layers of barrier materials are substantially visible to the observer at the same time identifying information may be displayed over the multi-layer barrier coating after it is applied to the plastic tube.

The first layer of the multi-layer barrier coating may be formed on the tube by dip-coating, roll-coating or spraying

an aqueous emulsion of the acrylate polymer, followed by air drying.

The acrylate polymer material may also be applied to the tube by an evaporation and curing process carried out as described in U.S. Patent No. 5,032,461, the disclosure of which is herein incorporated by reference.

5 The acrylate evaporation and curing process involves first atomizing the acrylate monomer into about 50 micron droplets and then flashing them off of a heated surface. This produces an acrylate molecular vapor which has the same chemistry as the starting monomer.

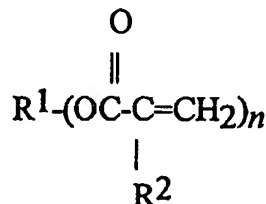
Acrylates are available with almost any chemistry desired. They usually have either one, two or three acrylate groups per molecule. Various mixtures of mono, di and tri acrylates are useful in the present invention. Most preferable are monoacrylates and diacrylates.

10 Acrylates form one of the most reactive classes of chemicals. They cure rapidly when exposed to UV or electron beam radiation to form a cross-linked structure. This imparts high temperature and abrasion resistant properties in the coating.

15 The monomer materials utilized are relatively low in molecular weight, between 150 and 1,000 and preferably in the range of 200 to 300 and have vapor pressures between about  $1 \times 10^{-6}$  Torr and  $1 \times 10^{-1}$  Torr at standard temperature and pressure (i.e., relatively low boiling materials). A vapor pressure of about  $1 \times 10^{-2}$  Torr is preferred. Polyfunctional acrylates are especially preferred. The monomers employed have at least two double bonds (i.e., a plurality of olefinic groups). The high-vapor-pressure monomers used in the present invention can be vaporized at low temperatures and thus are not degraded (cracked) by the heating process. The absence of unreactive degradation products means that films formed from these low molecular weight, high-vapor-pressure monomers have reduced volatile levels of components.

20 As a result, substantially all of the deposited monomer is reactive and will cure to form an integral film when exposed to a source of radiation. These properties make it possible to provide substantially continuous coating despite the fact that the film is very thin. The cured film exhibits excellent adhesion and are resistant to chemical attack by organic solvents and inorganic salts.

25 Because of their reactivity, physical properties and the properties of cured films formed from such components, polyfunctional acrylates are particularly useful monomeric materials. The general formula for such polyfunctional acrylates is:



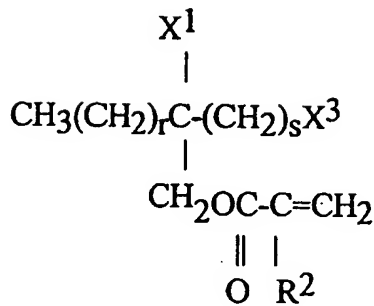
30 wherein:

40  $\text{R}^1$  is an aliphatic, alicyclic or mixed aliphatic-alicyclic radical;  
 $\text{R}^2$  is a hydrogen, methyl, ethyl, propyl, butyl or pentyl; and  
 $n$  is from 2 to 4.

45 Such polyfunctional acrylates may also be used in combination with various monacrylates, such as those having the formula:

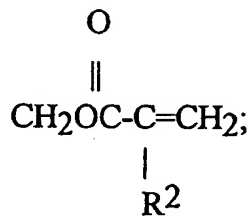
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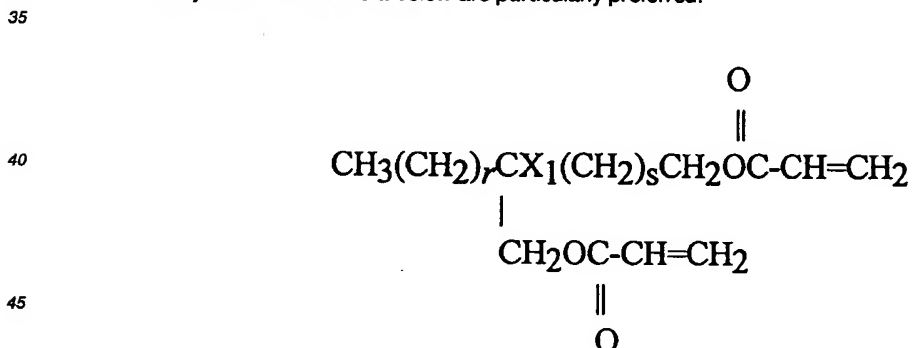


15 wherein:

20  $\text{R}^2$ , is as defined above;  
 $X^1$  is H, epoxy, 1,6-hexanediol, tripropylene glycol or urethane; and  
 $r, s$  are 1-18.



and  
 $X^3$  is CN or COOR<sup>3</sup> wherein R<sup>3</sup> is an alkyl radical containing 1-4 carbon atoms. Most often,  $X^3$  is CN or COOCH<sub>3</sub>.  
 Diacrylates of the formula below are particularly preferred:



50 wherein:  
 $X_1, r$  and  $s$  are as defined above.

Curing is accomplished by opening the double bonds of the reactant molecules. This can be accomplished by means of an energy source such as apparatus which emits infrared, electrons or ultraviolet radiation.

FIG. 6 illustrates the process for applying an acrylate coating. An acrylate monomer 100 is directed through a dielectric evaporator 102 and then through an ultrasonic atomizer 104 and into a vacuum chamber 106. The monomer droplets are atomized ultrasonically and the droplets vaporized where they condense on the tube or film that is loaded on a drum 108.

The condensed monomer liquid subsequently is radiation cured by means of an electron beam gun 110.

The second layer of the multi-layer barrier coating, an inorganic material, may be formed over the acrylate coating

by radio frequency discharge, direct or dual ion beam deposition, sputtering or plasma chemical vapor deposition, as described in U.S. Patent Nos. 4,698,256, 4,809,876, 4,992,298 and 5,055,318, the disclosures of which are herein incorporated by reference.

For example, a method of depositing an oxide coating is provided by establishing a glow discharge plasma in a previously evacuated chamber. The plasma is derived from one or more of the gas stream components, and preferably is derived from the gas stream itself. The article is positioned in the plasma, preferably adjacent the confined plasma, and the gas stream is controllably flowed into the plasma. A silicon oxide based film is deposited on the substrate to a desired thickness. The thickness of the oxide coating is about 100 Angstroms ( $\text{\AA}$ ) to about 10,000  $\text{\AA}$ . A thickness of less than about 10,000  $\text{\AA}$  may not provide sufficient barrier and a thickness of greater than about 10,000  $\text{\AA}$  may crack, thus decreasing the effective barrier. Most preferably, the thickness of the oxide coating is about 1,000  $\text{\AA}$  to about 3,000  $\text{\AA}$ .

Another method of depositing an oxide coating is by confining a plasma with magnets. Preferably, the magnetically enhanced method for depositing a silicon oxide based film on a substrate is preferably conducted in a previously evacuated chamber of glow discharge from a gas stream. The gas stream preferably comprises at least two components: a volatilized organosilicon component, an oxidizer component such as oxygen, nitrous oxide, carbon dioxide or air and an optionally inert gas component.

Examples of suitable organosilicon compounds useful for the gas stream in the plasma deposition methods are liquid or gas at about ambient temperature and when volatilized have a boiling point about 0°C to about 150°C and include dimethylsilane, trimethylsilane, diethylsilane, propylsilane, phenylsilane, hexamethyldisilane, 1,1,2,2-tetramethyldisilane, bis(trimethylsilane)methane, bis(dimethylsilyl) methane, hexamethyldisiloxane, vinyl trimethoxy silane, vinyl triethoxysilane, ethylmethoxysilane, ethyltrimethoxysilane, divinyltetramethyldisiloxane, hexamethyldisilazane divinylhexamethyltrisiloxane, trivinylpentamethyltrisiloxane, tetraethoxysilane and tetramethoxysilane.

Among the preferred organosilicons are 1,1,3,3-tetramethyldisiloxane, trimethylsilane, hexamethyldisiloxane, vinyltrimethylsilane, methyltrimethoxysilane, vinyltrimethoxysilane and hexamethyldisilazane. These preferred organosilicon compounds have boiling points of 71°C, 55.5°C, 102°C, 123°C and 127°C respectively.

The optional inert gas of the gas stream preferably is helium, argon or nitrogen.

The volatilized organosilicon component is preferably admixed with the oxygen component and the inert gas component before being flowed into the chamber. The quantities of these gases being so admixed are controlled by flow controllers so as to adjustably control the flow rate ratio of the gas stream components.

Various optical methods known in the art may be used to determine the thickness of the deposited film while in the deposition chamber, or the film thickness can be determined after the article is removed from the deposition chamber.

The deposition method of the present invention is preferably practiced at relatively high power and quite low pressure. A pressure less than about 500 millitorr (mTorr) should be maintained during the deposition, and preferably the chamber is at a pressure between about 43 to about 490 millitorr during the deposition of film. Low system pressure results in lower deposition rates whereas higher system pressure provides faster deposition rates. When the plastic article to be coated is heat sensitive, a higher system pressure may be used to minimize the amount of heat the substrate is exposed to during deposition because high substrate temperatures are to be avoided for low T<sub>g</sub> polymers such as polypropylene and PET (T<sub>g</sub> is -10°C and 60°C respectively).

The substrate is electrically isolated from the deposition system (except for electrical contact with the plasma) and is at a temperature of less than about 80°C during the depositing. That is, the substrate is not deliberately heated.

Referring to FIG. 7, the system for depositing a silicon oxide based film comprises an enclosed reaction chamber 170 in which a plasma is formed and in which a substrate or tube 171, is placed for depositing a thin film of material on a sample holder 172. The substrate can be any vacuum compatible material, such as plastic. One or more gases are supplied to the reaction chamber by a gas supply system 173. An electric field is created by a power supply 174.

The reaction chamber can be of an appropriate type to perform any of the plasma-enhanced chemical vapor deposition (PECVD) or plasma polymerization process. Furthermore, the reaction chamber may be modified so that one or more articles may be coated with an oxide layer simultaneously within the chamber.

The pressure of the chamber is controlled by a mechanical pump 188 connected to chamber 170 by a valve 190.

The tube to be coated is first loaded into chamber 170 in sample holder 172. The pressure of the chamber is reduced to about 5m Torr by mechanical pump 188. The operating pressure of the chamber is about 90 to about 140 mTorr for a PECVD or plasma polymerization process and is achieved by flowing the process gases, oxygen and trimethyl silane, into the chamber through monomer inlet 176.

The thin film is deposited on the outer surface of the tube and has a desired uniform thickness or the deposition process may be interrupted periodically to minimize heating of the substrate and/or electrodes and/or physically remove particulate matter from the articles.

Magnets 196 and 198 are positioned behind electrode 200 to create an appropriate combination of magnetic and electrical fields in the plasma region around the tube.

The system is suitable for low frequency operation. An example frequency is 40kHz. However, there can be some advantages from operating at a much high frequency, such as in the radio frequency range of several megahertz.

The silicon oxide based film or blends thereof used in accordance with this disclosure, may contain conventional

additives and ingredients which do not adversely affect the properties of articles made therefrom.

A variety of substrates can be coated with a barrier coating by the process of the present invention. Such substrates include, but are not limited to packaging, containers, bottles, jars, tubes and medical devices.

5 A plastic blood collection tube coated with the multi-layer barrier coating will not interfere with testing and analysis that is typically performed on blood in a tube. Such tests include but are not limited to, routine chemical analysis, biological inertness, hematology, blood chemistry, blood typing, toxicology analysis or therapeutic drug monitoring and other clinical tests involving body fluids. Furthermore, a plastic blood collection tube coated with the barrier coating is capable of being subjected to automated machinery such as centrifuges and may be exposed to certain levels of radiation in the sterilization process with substantially no change in optical or mechanical and functional properties.

10 A plastic blood collection tube coated with the multi-layer barrier coating is able to maintain 90% original draw volume over a period of one year. Draw volume retention depends on the existence of a particle vacuum, or reduced pressure, inside the tube. The draw volume changes in direct proportion to the change in vacuum (reduced pressure). Therefore, draw volume retention is dependent on good vacuum retention. A plastic tube coated with a barrier coating substantially prevents gas permeation through the tube material so as to maintain and enhance the vacuum retention  
15 and draw volume retention of the tube. Plastic tubes without the multi-layer coating of the present invention may maintain about 90% draw volume for about 3 to 4 months.

If the multi-layer barrier coating is also coated or applied on the inner surface of the plastic blood collection tube, the barrier coating may be hemorepellent and/or have characteristics of a clot activator.

It will be understood that it makes no difference whether the plastic composite container is evacuated or not evacuated in accordance with this invention. The presence of a barrier coating on the outer surface of the container has the effect of maintaining the general integrity of the container holding a sample so that it may be properly disposed of without any contamination to the user. Notable is the clarity of the barrier coating as coated or applied on the container and its abrasion and scratch resistance.

The barrier coating used in accordance with this disclosure, may contain conventional additives and ingredients  
25 which do not adversely affect the properties of articles made therefrom.

The following examples are not limited to any specific embodiment of the invention, but are only exemplary.

#### EXAMPLE 1

##### 30 METHOD FOR COATING PLASTIC SUBSTRATES TUBES WITH MULTI-LAYER BARRIER COATING

An acrylate coating was applied to polypropylene tubes and films (substrates) of various thickness in a chamber wherein Tripropylene Glycol Diacrylate (TPGDA) was fed into the evaporator and was flash vaporized at about 343°C onto the substrate in the chamber and condensed. The condensed monomer film was then E-beam cured by an electron beam gun.  
35

The substrate coated with the acrylate coating (TPGDA) was then cleaned with a mixture comprising equal parts of a micro detergent and de-ionized (DI) water solution. The substrate was rinsed thoroughly in DI water and allowed to air dry. The cleaned substrate was then stored in a vacuum oven at room temperature until it was to be coated.

40 The cleaned substrate was then attached to a holder which fits midway between the electrodes in the glass vacuum chamber. The chamber was closed and a mechanical pump was used to achieve a base pressure of 5 mTorr.

The electrode configuration is internally capacitively coupled with permanent magnets on the backside of the titanium electrodes. This special configuration provides the ability to confine the glow between the electrodes because of the increase in collision probability between electrons and reacting gas molecules. The net result of applying a magnetic field is similar to increasing the power applied to the electrodes, but without the disadvantages of higher bombardment energies and increased substrate heating. The use of magnetron discharge allows operation in the low pressure region and a substantial increase in polymer deposition rate.  
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The monomer which consists of a mixture of trimethylsilane (TMS) and oxygen was introduced through stainless steel tubing near the electrodes. The gases were mixed in the monomer inlet line before introduction into the chamber. Flow rates were manually controlled by stainless steel metering valves. A power supply operating at an audio frequency  
50 of 40 kHz was used to supply power to the electrodes. The system parameters used for thin film deposition of plasma polymerized TMS/O<sub>2</sub> on the polymer substrate were as follows:

5	Surface Pretreatment:	TMS Flow Base Pressure Oxygen Flow System Pressure Power Time	= 0 sccm = 5 mTorr = 10 sccm = 140 mTorr = 50 watts = 2 minutes
10	Oxide Deposition:	TMS Flow Oxygen Flow System Pressure Power Deposition Time	= 0.75 - 1.0 sccm = 2.5 = 3.0 sccm = 90 - 100 mTorr = 30 watts = 5 minutes
15			
20			

After the thin film was deposited, the reactor was allowed to cool. The reactor was then opened, and the substrate with a multi-layer barrier coating was removed.

#### EXAMPLE 2

#### **METHOD FOR COATING PLASTIC SUBSTRATES WITH MULTI-LAYER BARRIER COATING**

An acrylate coating was applied to polypropylene tubes and films (substrates) in a chamber wherein a 60:40 mixture of isobornyl acrylate: epoxydiacrylate (IBA:EDA) was fed into the evaporator and flash vaporized at about 343°C onto the substrate in the chamber and condensed. The condensed monomer film was then UV cured by an actinic light source of 365 nm.

The substrate coated with the acrylate coating (IBA:EDA) was then cleaned with a mixture comprising equal parts of a micro detergent and de-ionized (DI) water solution. The substrate was rinsed thoroughly in DI water and allowed to air dry. The cleaned substrate was then stored in a vacuum oven at room temperature until it was to be coated.

The cleaned substrate was then attached to a holder which fits midway between the electrodes in the glass vacuum chamber. The chamber was closed and a mechanical pump was used to achieve a base pressure of 5 mTorr.

The electrode configuration is internally capacitively coupled with permanent magnets on the backside of the titanium electrodes. The special configuration provides the ability to confine the glow between the electrodes because of the increase in collision probability between electrons and reacting gas molecules. The net result of applying a magnetic field is similar to increasing the power applied to the electrodes, but without the disadvantages of higher bombardment energies and increased substrate heating. The use of magnetron discharge allows operation in the low pressure region and a substantial increase in polymer deposition rate.

The monomer which consists of a mixture of trimethylsilane (TMS) and oxygen was introduced through stainless steel tubing near the electrodes. The gases were mixed in the monomer inlet line before introduction into the chamber. Flow rates were manually controlled by stainless steel metering valves. A power supply operating at an audio frequency of 40 kHz was used to supply power to the electrodes. The system parameters used for thin film deposition of plasma polymerized TMS/O<sub>2</sub> on the polymer substrate were as follows:

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5	Surface Pretreatment:	TMS Flow Base Pressure Oxygen Flow System Pressure Power Time	= 0 sccm = 5 mTorr = 10 sccm = 140 mTorr = 50 watts = 2 minutes
10	Oxide Deposition:	TMS Flow Oxygen Flow System Pressure Power Deposition Time	= 0.75 - 1.0 sccm = 2.5 - 3.0 sccm = 90 - 100 mTorr = 30 watts = 5 minutes
15			

20 After the thin film was deposited, the reactor was allowed to cool. The reactor was then opened, and the substrate with a multi-layer barrier coating was removed.

### EXAMPLE 3

#### 25 COMPARISON OF SUBSTRATES WITH AND WITHOUT MULTI-LAYER BARRIER COATINGS

All of the substrates prepared in accordance with Examples 1 and 2 above were evaluated for oxygen permeance (OTR) in the oxide coatings.

##### 30 (i) Oxygen permeance (OTR):

35 Film or plaque samples were tested for oxygen permeance (OTR) using a MO CON Ox-TRAN 2/20 (sold by Modern Controls, Inc., 7500 Boone Avenue N., Minneapolis, MN 55428). A single side of the film sample was exposed to 1 atm of 100% oxygen atmosphere. Oxygen permeating through the sample film was entrained in a nitrogen carrier gas stream on the opposite side of the film, and detected by a coulometric sensor. An electrical signal was produced in proportion to the amount of oxygen permeating through the sample. Samples were tested at 30°C and 0% relative humidity (R.H.). Samples were conditioned for 1 to 20 hours prior to determining oxygen permeance. The results are reported in Table 1 in units of cc/m<sup>2</sup>-atm-day.

40 Tube samples were tested for oxygen permeance (OTR) using a MOCON O<sub>x</sub>-TRAN 1,000 (sold by Modern Controls, Inc., 7500 Boone Avenue N., Minneapolis, MN 55428). A package adapter was used for mounting the tubes in a manner that allowed the outside of the tube to be immersed in a 100% O<sub>2</sub> atmosphere while the inside of tube is flushed with a nitrogen carrier gas. The tubes were then tested at 20°C and 50% R.H. The tubes were allowed to equilibrate for 2-14 days before a steady state permeability is determined. The results are reported in Table 1 in units of cc/m<sup>2</sup>-atm-day.

##### 45 (ii) Atom % of the elements present in the oxide coating:

A Surface Science Model SSx-100 X-ray photoelectron spectrometer (ESCA) was used to determine the atom % of the elements present in the oxide coatings. Film samples were placed inside the spectrometer and the elemental composition was determined of approximately 100Å into the surface. The surface was then argon ion etched as follows: 5000V and 9-10 mA argon ion beam was directed at the sample surface. After 5 seconds the ESCA spectra was taken and this procedure was repeated for a total of 5 times. The etch time was then increased to 20 seconds followed by ESCA and this process was repeated for a total of ten times. Finally, the etch time was increased to 40 second and the ESCA spectra was obtained until the bulk acrylate or the polymer substrate was reached. The oxide layer was clearly indicated by the presence of silicon in the ESCA spectra between the etch times of about 0 through about 1.3 minutes.

## (iii) Analysis of the Data in Table 1 and Figures 8 and 9:

Figures 8 and 9 illustrate atom % verses etch time for oxide samples deposited on polypropylene with acrylate primer coating and without acrylate coating. The data in Table 1 demonstrates for these plasma deposition processes the improved barrier properties correlate well with improved gas-barrier performance as a result of using the multi-layer barrier coating. The SiO<sub>x</sub> deposition on an acrylate treated surface occurs with less surface degradation and therefore resulting in a more dense barrier structure. Since the acrylate coating itself provides no detectable improvement in barrier, data demonstrates that the acrylate-oxide coating method greatly improves the barrier properties of plastic substrates.

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TABLE 1

Sample	Acrylate Coating	SiO <sub>x</sub> Coating Method	Oxygen Transmission (cc/m <sup>2</sup> -atm-da)
PP film, control	no	no	1500
PP plaque, 75 mil	no	yes	37
PP plaque, 75 mil	TPGDA	yes	3.5
PP plaque, 75 mil	no	yes	65
PP tube, control	no	no	70
PP tube	IBA:DA	no	70
PP tube	no	yes	46-60
PP tube	IBA:DA	yes	4.3-6.4

IBA:DA = iso-norbornyl: epoxydiacrylate (60:40), UV cured  
 TPGDA = tripropylene glycol diacrylate E-beam cured  
 Oxide Coatings = 1000 - 3000 Angstroms (as measured by Scanning Electron Microscope)  
 PP = polypropylene  
 plaque = 75 mil thickness  
 film = 2 mil thickness  
 tubes = nominal wall thickness of 40 mil

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## Claims

40 1. A sample assembly comprising:  
 a plastic container comprising polyethylene terephthalate, polypropylene or polyethylene napthalate, having an open end, a closed end, an inner surface and an outer surface; and  
 45 a multi-layer barrier coating associated over the outer surface of said container, said coating having a first layer comprising an acrylate primer coating material and a second layer adjacent on said first layer comprising a silicon oxide or an aluminum oxide.

2. The assembly of Claim 1, further comprising a closure in said open end of said container whereby a container and closure interface is formed.

50 3. The assembly of Claim 2, wherein said plastic container is a tube and said closure is an elastomeric stopper.

4. The assembly of Claim 2, wherein said tube is evacuated.

55 5. The assembly of Claim 1, wherein said multi-layer barrier coating includes registered tamper serrations adjacent to said container and said closure interface.

6. The assembly of Claim 1, wherein said first layer is a polymerized blend of mono-and di-acrylates.

7. The assembly of Claim 1, wherein said second layer is deposited by radio-frequency discharge, direct ion beam deposition, dual ion beam deposition, sputtering, plasma chemical vapor deposition or magnetically enhanced plasma systems.

5       8. The assembly of Claim 1, further comprising a multi-layer barrier coating over the inner surface of said container, having a first layer including an acrylate primer coating material and a second layer on said first layer of a metal oxide.

10      9. The assembly of Claim 1, wherein said first layer is deposited on said outer surface of said container in a previously evacuated chamber comprising the following steps:

15           (a) selecting a curable component comprising: i) polyfunctional acrylates, or ii) mixtures of monoacrylates and polyfunctional acrylates;

             (b) flash vaporizing said component into said chamber;

             (c) condensing a film of vaporized component onto the outer surface of said container; and

             (d) curing said film.

10. The assembly of Claim 9, wherein said second layer is deposited in said previously evacuated chamber onto said first layer by the following steps:

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FIG-1

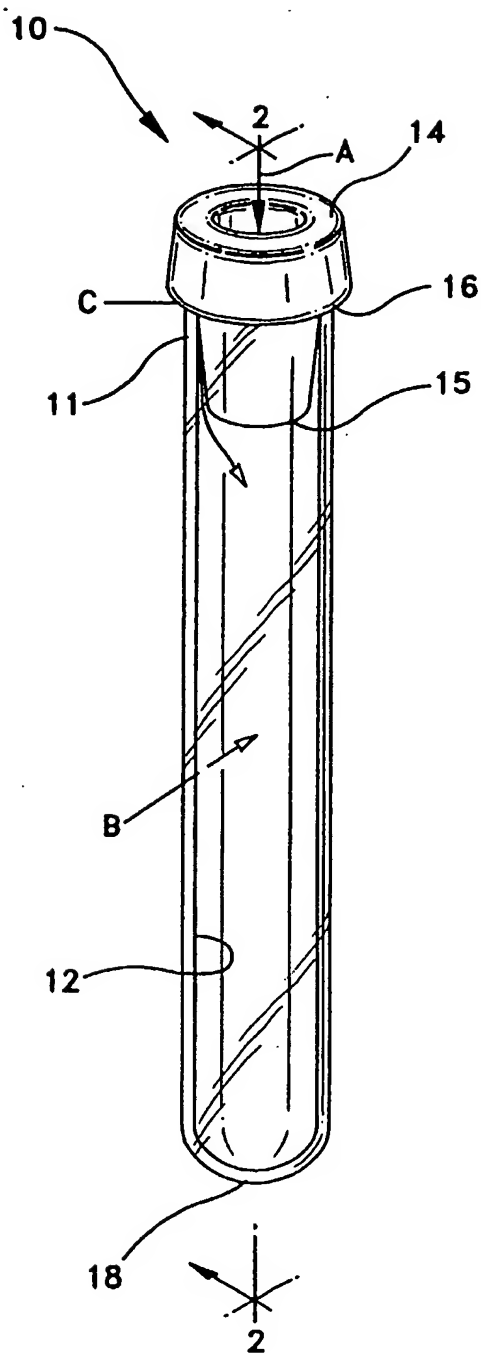


FIG-2

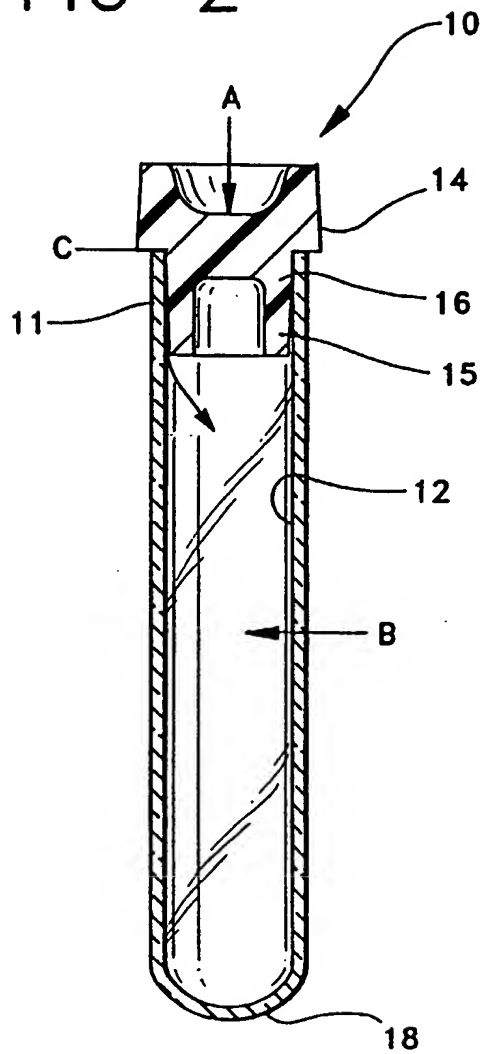


FIG-3

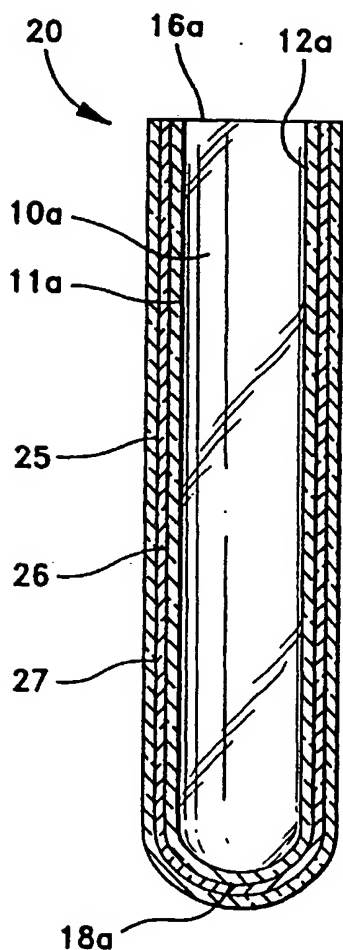


FIG-4

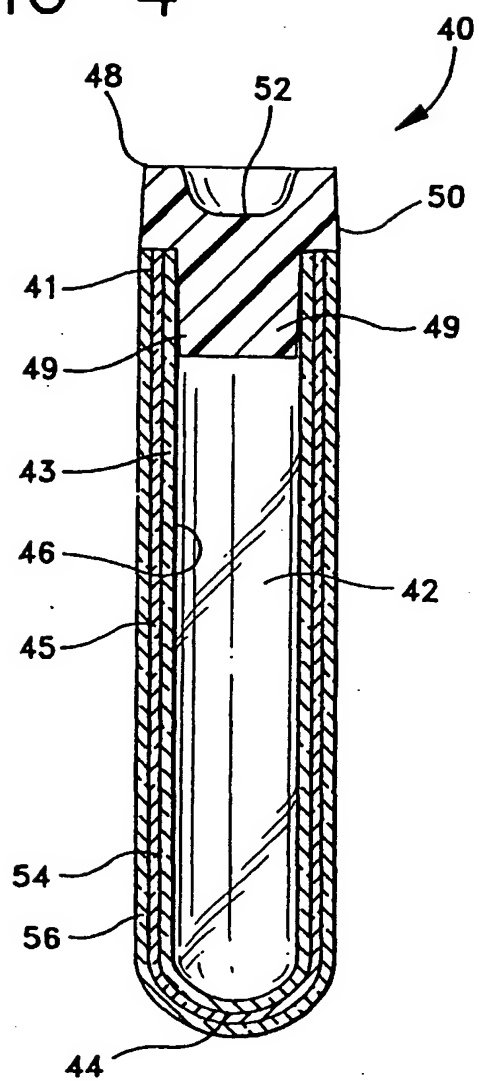


FIG-5

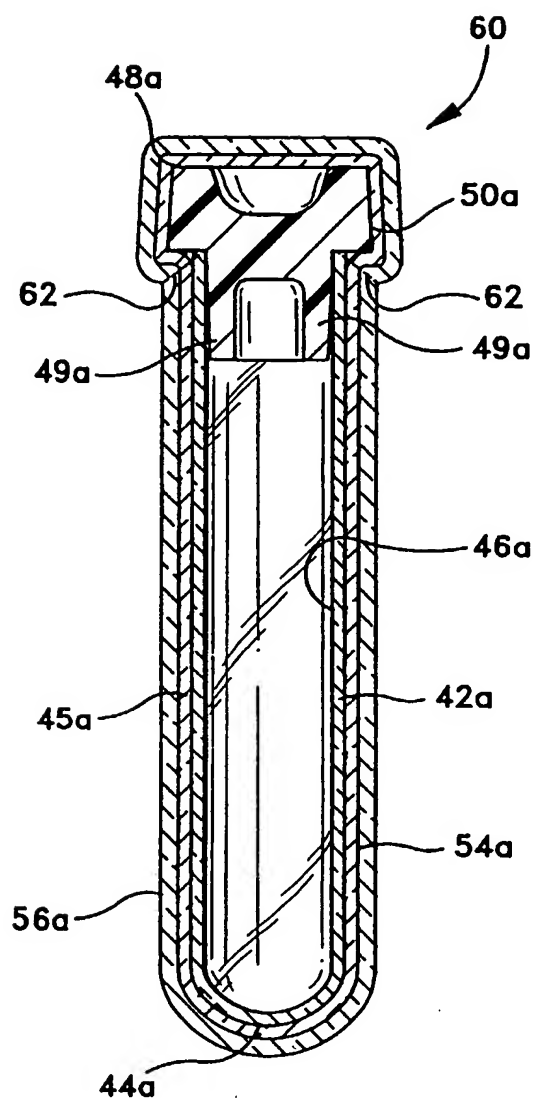


FIG-6

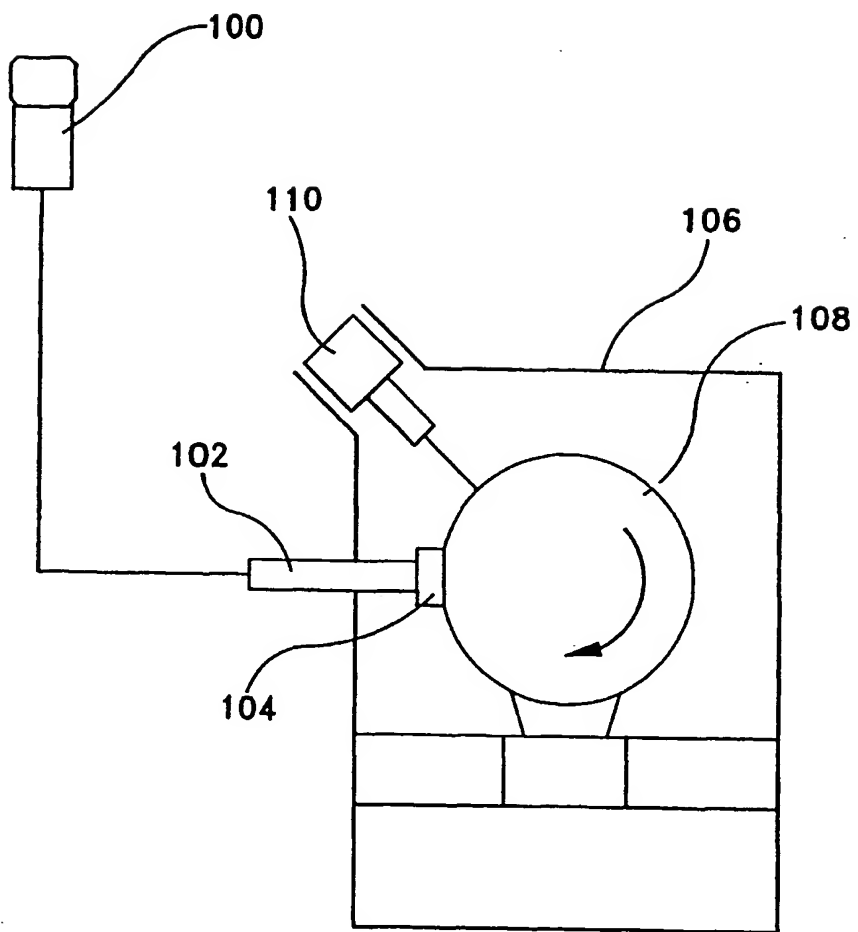


FIG-7

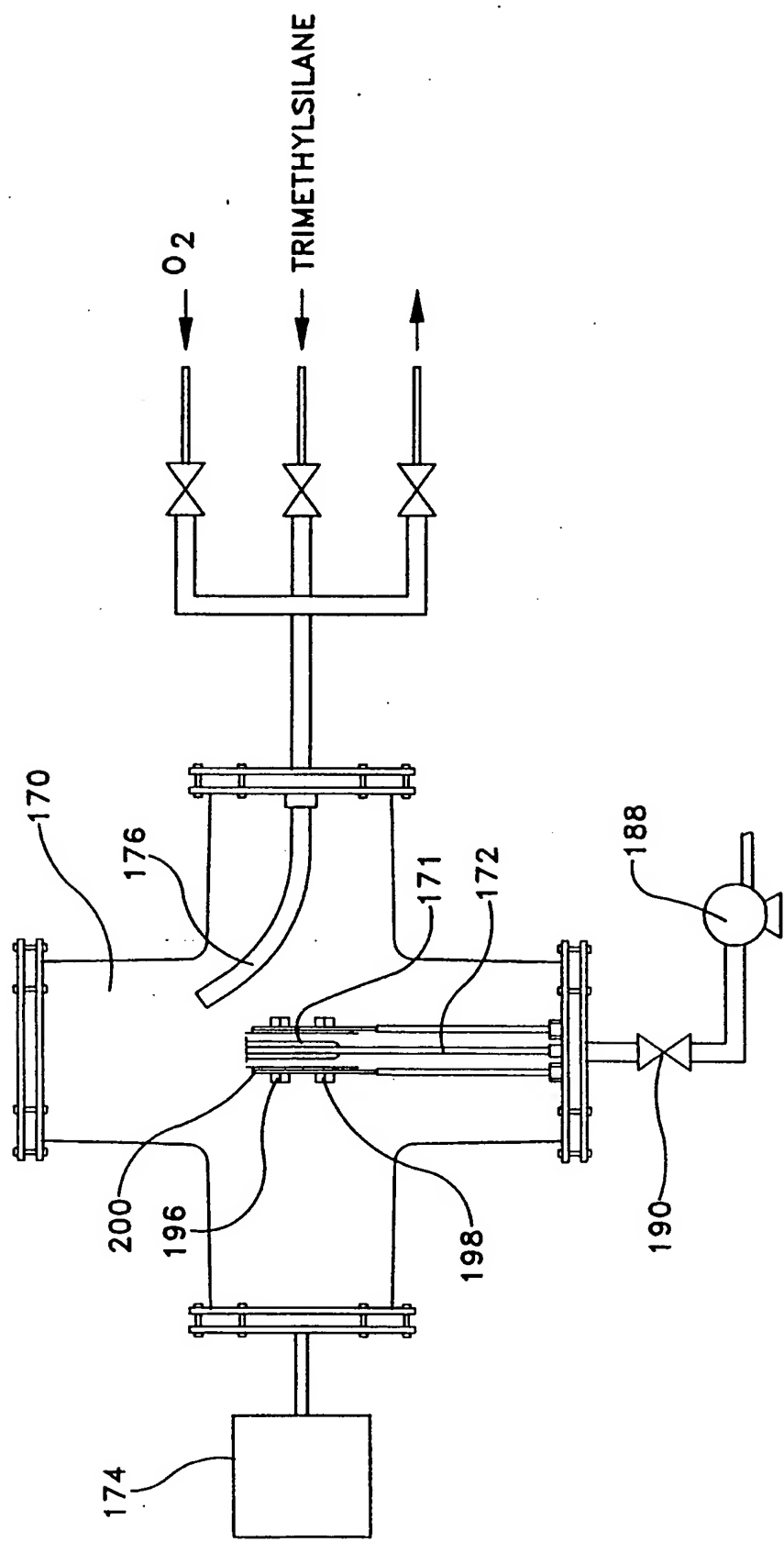


FIG-8

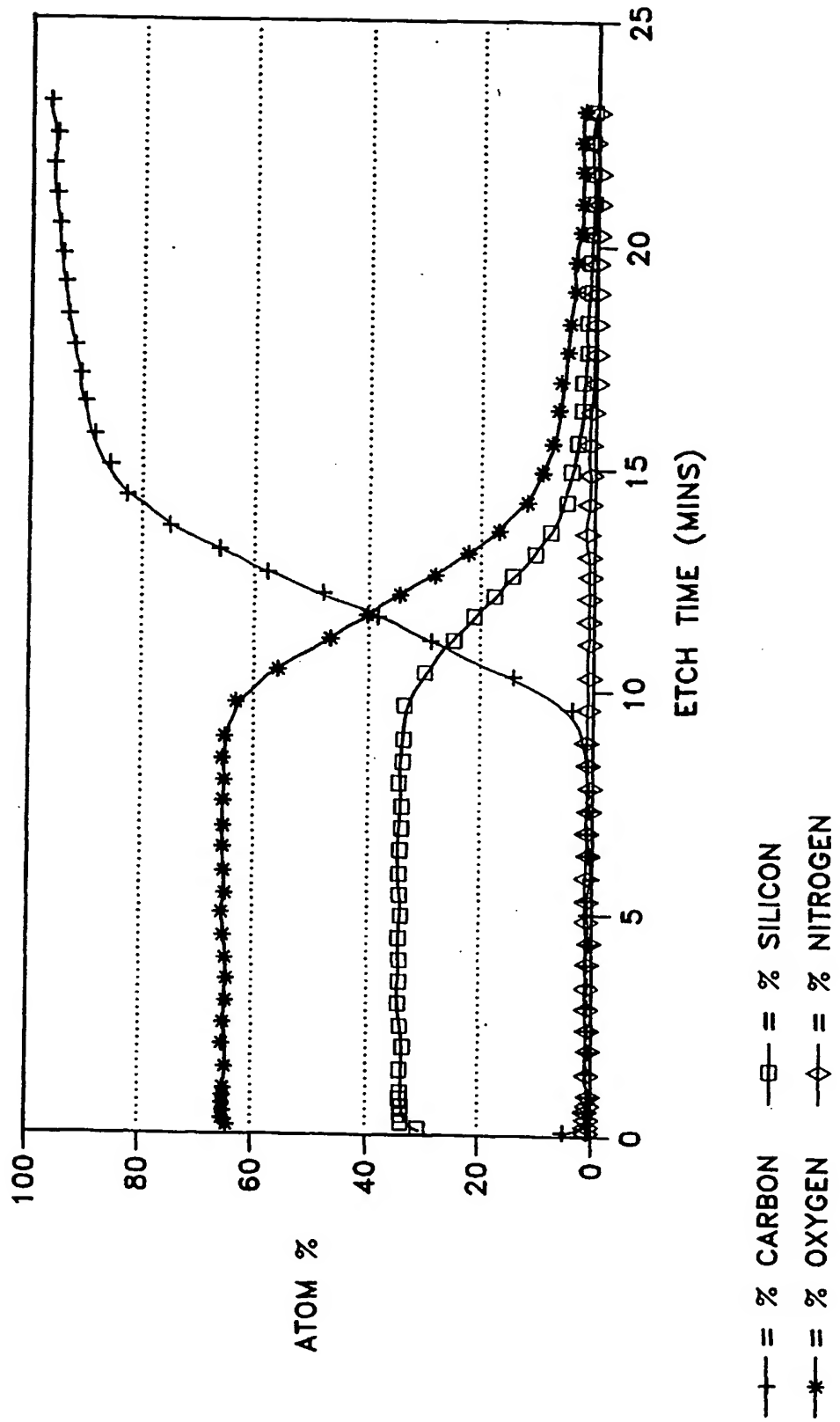
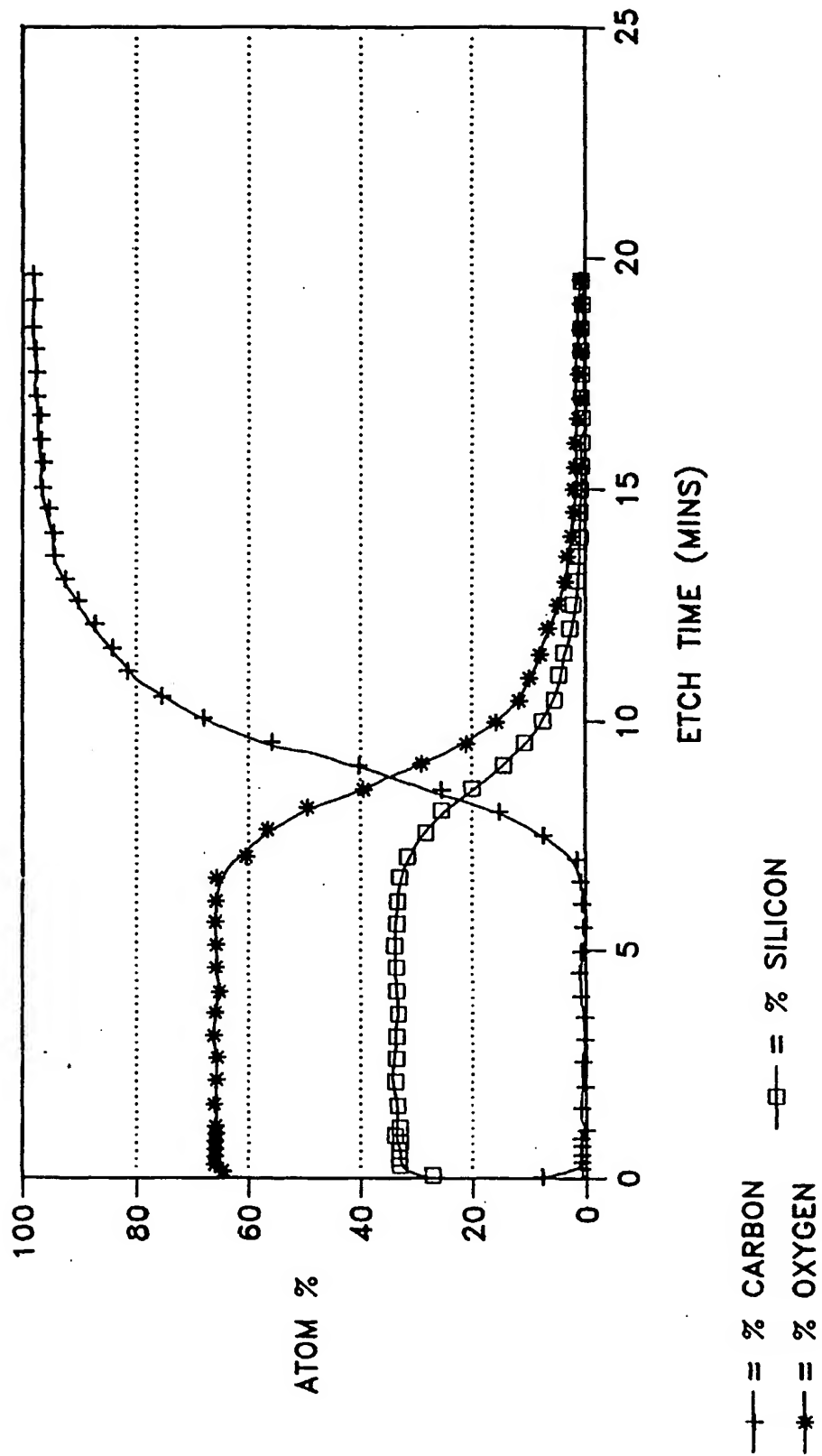


FIG-9





European Patent  
Office

### PARTIAL EUROPEAN SEARCH REPORT

Application Number

which under Rule 45 of the European Patent Convention EP 96 10 3454  
shall be considered, for the purposes of subsequent  
proceedings, as the European search report

DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
X	US-A-5 440 446 (SHAW DAVID G ET AL) 8 August 1995	1,6,7,9	B01L3/14 B32B27/00 C23C14/00
Y	* column 5, line 9 - column 7, line 38 * * column 16, line 53 - column 18, line 63 *	2-5,8	
Y	---	2-5	
Y	EP-A-0 535 810 (BECTON DICKINSON CO) 7 April 1993 * page 3, line 56 - page 4, line 41 *	---	
Y	EP-A-0 603 717 (BECTON DICKINSON CO) 29 June 1994 * page 4, line 50 - line 52 *	8	
D,A	US-A-4 842 893 (YIALIZIS ANGELO ET AL) 27 June 1989 * column 2, line 13 - column 3, line 35 *	---	
A	US-A-4 200 681 (HALL WALTER L ET AL) 29 April 1980 * the whole document *	-----	
			TECHNICAL FIELDS SEARCHED (Int.Cl.6)  B01L B32B C23C
<b>INCOMPLETE SEARCH</b>			
<p>The Search Division considers that the present European patent application does not comply with the provisions of the European Patent Convention to such an extent that it is not possible to carry out a meaningful search into the state of the art on the basis of some of the claims</p> <p>Claims searched completely :          Claims searched incompletely :          Claims not searched :          Reason for the limitation of the search:</p> <p>see sheet C</p>			
Place of search	Date of completion of the search	Examiner	
THE HAGUE	5 August 1996	Bindon, C	
CATEGORY OF CITED DOCUMENTS		T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document	
<small>EPO FORM 150 012 (PACOM)</small>			

**EP 0 794 007 A1**



**EP 96 10 3454**

**-C-**

**INCOMPLETE SEARCH**

Claims searched completely	:	1-9
Claim not searched	:	10

**Reason: Claim 10 is clearly incomplete**

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